## **Amendments to the Claims**

This listing of claims will replace all prior versions, and listings, of the claims in the application.

## Listing of Claims

- 1. (Currently amended) A pharmaceutical composition comprising
  - a biologically active agent; (i)
  - (ii) an a first adjuvant chemical which increases the effect of the biologically active agent, said chemical selected from one or more of:
    - A) a polyamino acid polyornithine,
    - B) a water soluble vitamin or water soluble vitamin derivative,
    - C) a positively charged cationic pluronics block copolymer or a positively charged cationic surfactant,
    - D) a clathrate,
    - E) a complexing agent,
    - cetrimides, F)
    - G) an S-layer protein, or
    - H) Methyl-glucamine; and
  - (iii) a pharmaceutically acceptable carrier or diluent; subject to the following provisos

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a) when the chemical (ii) above is selected from D) or E), the biologically

active agent is an agent which is capable of generating that generates a

protective immune response in an animal to which it is administered;

b) when the chemical (ii) above is selected from A) and the biologically active

agent is an agent which is capable of generating that generates a protective

immune response in an animal to which it is administered, the composition is .

for administration to a mucosal surface,

c) when the chemical (ii) above is selected from C) and the biologically active

agent is an agent which is capable of generating that generates a protective

immune response in an animal to which it is administered, the composition

does not contain a polyacrylic acid, and

d) when the chemical (ii) above is selected from G) and the biologically active

agent is an agent which is capable of generating that generates a protective

immune response in an animal to which it is administered, the carrier or

diluent of (iii) is a microsphere or liposome.

2. (Currently amended) A The composition according to of claim 1

wherein the biologically active agent is an agent that is capable of generating generates a

protective immune response in an animal to which it is administered.

- 3. (Currently amended) A The composition according to of claim 1 wherein the said adjuvant chemical can act acts as an immunostimulant.
- 4. (Currently amended) A The composition according to of claim 1 wherein the said adjuvant chemical is selected from one or more of;
  - A) the poly-ornithine has a, for example of molecular weight from 5 to 150kDa;
- B) the water soluble vitamin vitamins or water soluble vitamin derivative derivatives is such as vitamin E TPGS (d-alpha tocophenyl polyethylene glycol 1000 succinate),
- C) <u>the cationic pluronics which are block copolymer copolymers or the cationic surfactant is surfactants which are positively charged by means of, in particular with NH<sub>2</sub><sup>+</sup> groups</u>
- D) the complexing agent forms agents which form complexes with fatty acids such as deoxycholic acid, or
- E) the clathrate is a cyclodextrin or a derivative thereof eyelodextrins and their derivatives such as dimethyl  $\beta$  cyclodextrin.
- 5. (Currently amended) A The composition according to of claim 1 wherein the carrier comprises a particle.

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- 6. (Currently amended) A The composition according to of claim 5 wherein the particle is a microsphere or liposome.
- 7. (Currently amended) A The composition according to of claim 6 which comprises a microsphere.
- 8. (Currently amended) A The composition according to of claim 7 wherein the microsphere is prepared using a high molecular weight polymer.
- 9. (Currently amended) A The composition according to of claim 8 wherein the polymer has a molecular weight of 100kDa or more.
- 10. (Currently amended) A <u>The</u> composition according to <u>of</u> claim 7 wherein the microsphere comprises poly-(L-lactide).
- 11. (Currently amended) A The composition according to of claim 1 wherein the ratio of the chemical (ii) to the carrier (iii) is from 99:1 to 9:1 w/w.
- 12. (Currently amended) A The composition according to of claim 1 which is adapted for administration to a mucosal surface or is suitable for parenteral administration administered to a mucosal surface of the animal or administered parenterally to the animal.

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13. (Currently amended) A <u>The</u> composition according to <u>of</u> claim 2 which

further comprises a further-second adjuvant.

14. (Withdrawn) A method of producing a prophylactic or therapeutic

vaccine, which method comprises encapsulating a polypeptide which is capable of producing

a protective immune response in a first polymeric material which has a high molecular

weight, in the presence of a second polymeric material which increases the biological effect

of the composition.

15. (Withdrawn) A method of protecting a mammal against infection,

which method comprises administration of a composition according to claim 1 to a mammal.

16. (Withdrawn) A method according to claim 15 wherein the

composition is applied to a mucosal surface.

17. (Withdrawn) A method according to claim 16 wherein the mucosal

surface comprises an intranasal surface.

18. (Withdrawn) A microsphere comprising a polymeric carrier and an S-

layer protein.

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19. (Withdrawn) A microsphere according to claim 18 wherein said S-layer protein is coated on the surface of the microsphere.

20. (Withdrawn) A microsphere according to claim 18 which further comprises an agent that is capable of generating a protective immune response in an animal to which it is administered.

- 21. (Withdrawn) A microsphere according to claim 20 wherein one or more of said agents are linked to the S-layer protein.
- 22. (Withdrawn) A pharmaceutical composition comprising a microsphere according to claim 19.
- 23. (Withdrawn) A pharmaceutical composition according to claim 22 wherein said composition is a vaccine, intended to produce a protective immune response against a bacterium, and said S-layer protein is derived from said bacterium.
  - 24. (Withdrawn) The use of a chemical selected from
    - A) a polyamino acid,
    - B) a water soluble vitamin or vitamin derivative,

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- C) positively charged cationic pluronics,
- D) a clathrate,
- E) a complexing agent,
- F) cetrimides,
- G) an S-layer protein, or
- H) Methyl-glucamine

as an immunostimulant, provided that in the case of A), the immunostimulant is applied to a mucosal surface, in the case of C, the compound is used in the absence of a polyacrylic acid.

- 25. (Withdrawn) The use of an adjuvant chemical selected from
  - A) a polyamino acid,
  - B) a water soluble vitamin or vitamin derivative,
  - C) positively charged cationic pluronics,
  - D) a clathrate,
  - E) a complexing agent,
  - F) cetrimides,
  - G) an S-layer protein, or
  - H) Methyl-glucamine

as an immunostimulant in the production of a vaccine for use in prophylactic or therapeutic treatment, provided that in the case of A), the immunostimulant is used in a vaccine which is

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applied to a mucosal surface, in the case of C), the compound is used in the absence of a polyacrylic acid.

- 26. (New) The composition of claim 4 wherein
  - A) the complexing agent forms complexes with deoxycholic acid; or
  - B) the clathrate is dimethyl- $\beta$ -cyclodextrin.